```
SEQ ID NO: 38
RESULT 7
AAV88429
ID
     AAV88429 standard; cDNA; 481 BP.
XX
AC
     AAV88429;
XX
DT
     12-FEB-1999 (first entry)
XX
DE
     EST clone EM42.
XX
KW
     Expressed sequence tag; secreted protein; haematopoiesis regulator;
KW
     tissue growth; activin; inhibin; tumour invasion suppressor; EST; human;
KW
     chemotaxis; chemokinesis; haemostasis; gene therapy; thrombolysis;
KW
     receptor; ligand; anti-inflammatory; tumour inhibitor; ds.
XX
os
     Homo sapiens.
XX
PN
     US2002/0173635.
XX
PD
     21 NOV 2002.
XX
PF
     07 June 2002.
XX
PR
     10-APR-1997;
                    97US-0837312.
XX
PA
     (GEMY ) GENETICS INST INC.
XX
PI
     Agostino MJ, Jacobs K,
                             Lavallie ER, McCoy JM,
PI
     Racie LA, Spaulding V, Treacy M;
XX
DR
     WPI; 1999-070078/06.
XX
PT
     New polynucleotides encoding human secreted proteins - derived from
PT
     e.g. human blood, kidney, foetal lung, placenta, testes, brain,
PT
     ovary, pituitary, retina and colon cDNA libraries
XX
PS
     Claim 1; Page 394; 641pp; English.
XX
CC
     The present sequence represents an expressed sequence tag (EST), and is
     a polynucleotide of the invention. The polynucleotides of the invention
CC
CC
     are all secreted EST sequences isolated from a variety of human tissue
CC
     sources. The EST sequences and proteins encoded by them are predicted to
CC
     have useful biological activities which would make them suitable for
CC
     treating, preventing or ameliorating medical conditions in humans and
CC
     animals, although no supporting data is given. Suggested activities
CC
     include nutritional activity, immune stimulating or suppressing activity,
CC
     haematopoiesis regulating activity, tissue growth activity,
CC
     activin/inhibin activity, chemotactic/chemokinetic activity, haemostatic
CC
     and thrombolytic activity, receptor/ligand activity, anti-inflammatory
CC
     activity, cadherin/tumour invasion suppressor activity, tumour inhibition
CC
     activity. The EST sequences are also stated to be useful for gene
CC
     therapy.
XX
SO
     Sequence 481 BP; 89 A; 127 C; 114 G; 151 T; 0 other;
                          35.7%; Score 452.4; DB 20; Length 481;
  Best Local Similarity 99.8%; Pred. No. 1.4e-102;
  Matches 453; Conservative
                                0; Mismatches
                                                   1:
                                                       Indels
```

0;

Qy	484	AACTGGTGCTGGTTTCACATCACGACAGCAACGGGATTCTTTTGCTTTGAAAATGTTGCA 543
Db	22	AACTGGTGCTGGTTTCACATCACGACAGCAACGGGATTCTTTTGCTTTGAAAATGTTGCA 81
Qy	544	GTCCACCTGTCCAACTTGATCTTCCGGACATTTGACTTGTTTCTGGTTATCCACCATCTC 603
Db	82	GTCCACCTGTCCAACTTGATCTTCCGGACATTTGACTTGTTTCTGGTTATCCACCATCTC 141
Qy	604	TTTGCCTTTCTTGGCTTGCTTGGTCAATCTCCAAGCTGGCCACTATCTAGCT 663
Db	142	TTTGCCTTCTTGGCTTGCTTGGCTGCTCAAGCTGGCCACTATCTAGCT 201
Qy	664	ATGACCACGTTGCTCCTGGAGATGAGCACGCCCTTTACCTGCGTTTCCTGGATGCTCTTA 723
Db	202	ATGACCACGTTGCTCCTGGAGATGAGCACGCCCTTTACCTGCGTTTCCTGGATGCTCTTA 261
Qy	724	AAGGCGGGCTGGTCCGAGTCTCTGTTTTGGAAGCTCAACCAGTGGCTGATGATTCACATG 783
Db	262	AAGGCGGGCTGGTCCGAGTCTCTGTTTTGGAAGCTCAACCAGTGGCTGATGATTCACATG 321
Qу	784	TTTCACTGCCGCATGGTTCTAACCTACCACATGTGGTGGGTG
Db	322	TTTCACTGCCGCATGGTTCTAACCTACCACATGTGGTGGTGTGTTTCTGGCACTGGGAC 381
Qy	844	GGCCTGGTCAGCAGCCTGTATCTGCCTCATTTGACACTGTTCCTTGTCGGACTGGCTCTG 903
Db	382	GGCCTGGTCAGCAGCCTGTATCTGCCTCATTTGACACTGTTCCTTGTCGGACTGGCTCTG 441
Qу	904	CTTACGCTAATCATTAATCCATATTGGACCCATA 937
Db	442	CTTACGCTAATCATTAATCCATATTGGACCCAGA 475